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Mutations in *CENPE* define a novel kinetochore-centromeric mechanism for microcephalic primordial dwarfism

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Abstract Defects in centrosome, centrosomal-associated and spindle-associated proteins are the most frequent cause of primary microcephaly (PM) and microcephalic primordial dwarfism (MPD) syndromes in humans. Mitotic progression and segregation defects, microtubule spindle abnormalities and impaired DNA damage-induced G2-M cell cycle checkpoint proficiency have been documented in cell lines from these patients. This suggests that impaired mitotic entry, progression and exit strongly contribute to PM and MPD. Considering the vast protein networks involved in coordinating this cell cycle stage, the list of potential target genes that could underlie novel developmental disorders is large. One such complex network, with a direct microtubule-mediated physical connection to the centrosome, is the kinetochore. This centromeric-associated

structure nucleates microtubule attachments onto mitotic chromosomes. Here, we described novel compound heterozygous variants in *CENPE* in two siblings who exhibit a profound MPD associated with developmental delay, simplified gyri and other isolated abnormalities. *CENPE* encodes centromere-associated protein E (CENP-E), a core kinetochore component functioning to mediate chromosome congression initially of misaligned chromosomes and in subsequent spindle microtubule capture during mitosis. Firstly, we present a comprehensive clinical description of these patients. Then, using patient cells we document abnormalities in spindle microtubule organization, mitotic progression and segregation, before modeling the cellular pathogenicity of these variants in an independent cell system. Our cellular analysis shows that a pathogenic defect in CENP-E, a kinetochore-core protein, largely phenocopies *PCNT*-mutated microcephalic osteodysplastic primordial dwarfism-type II patient cells. *PCNT* encodes

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a centrosome-associated protein. These results highlight a common underlying pathomechanism. Our findings provide the first evidence for a kinetochore-based route to MPD in humans.

Introduction

Microcephalic primordial dwarfism (MPD) is the collective term for a group of clinically overlapping syndromes whose underlying genetic basis has started to emerge within the last decade or so. Specific examples include Seckel syndrome, microcephalic primordial dwarfism-type II (MOPDii) and Meier-Gorlin Syndrome (Gorlin et al. 1975; Hall et al. 2004; Majewski et al. 1982b; Seckel 1960). For these specific disorders, causative defects have been identified in central components of the ATR-dependent DNA damage response (DDR), in a centrosome-associated protein and in components of the DNA replication licensing machinery, respectively (Bicknell et al. 2011a, b; Griffith et al. 2008; Guernsey et al. 2011; O'Driscoll et al. 2003; Ogi et al. 2012; Qvist et al. 2011; Rauch et al. 2008). Defects in DNA double strand break repair and other genome stability pathways can also cause MPD (Buck et al. 2006; Ijspeert et al. 2013; Murray et al. 2014; O'Driscoll et al. 2001; Shaheen et al. 2014; Shamseldin et al. 2012).

Primary microcephaly (PM) with and without overt short stature is commonly caused by defects in centrosome and spindle pole-associated proteins (Mahmood et al. 2011; Poirier et al. 2013; Thornton and Woods 2009). These PM and MPD-associated defects collectively underscore the requirement for optimal cell cycle progression and checkpoint control for normal human development, including growth and neurogenesis. Currently, mutations in genes encoding core centrosome and centrosome-associated proteins represent the most frequent causative defect identified so far in MPDs. The centrosome is an important microtubule-organizing centre essential for coordinating G2-M progression, normal mitotic cell division and even cilia formation. Interestingly, defects in the ATR-dependent DDR and in DNA replication licensing machinery are also associated with centrosome abnormalities, and vice versa, suggestive of a multifactorial but interconnected underlying pathobiology of MPDs (Alderton et al. 2004, 2006; Griffith et al. 2008; Stiff et al. 2013).

The centromeric kinetochore of mitotic chromosomes is the attachment site for spindle microtubules, ensuring their amphitelic attachments onto all of the chromosomes, thereby enabling their alignment at metaphase (Hori and Fukagawa 2012; Santaguida and Musacchio 2009). This facilitates accurate and timely segregation. Unattached kinetochores activate the mitotic spindle assembly

checkpoint (SAC) (Cleveland et al. 2003). SAC activation delays mitosis until the kinetochores of each chromosome have properly attached to spindle microtubules (Musacchio 2011; Musacchio and Salmon 2007). Congenital defects in the SAC gene *BUB1B* which encodes the SAC kinase BubR1 cause mosaic variegated aneuploidy (MVA): an MPD associated with aneuploidy and elevated cancer incidence (Hanks et al. 2004; Shinya Matsuura et al. 2006).

In contrast to the centrosome-spindle pole, pathogenic defects in core kinetochore components are currently notably under-represented as a cause of MPD. Using an exome sequencing strategy, we describe novel compound heterozygous variants in *CENPE* in two siblings characterized by a profound MPD with severe developmental delay, simplified gyri and various isolated abnormalities. Centromere-associated protein-E (CENP-E) is a large (>300 kD) kinetochore-associated kinesin-like motor protein required for spindle microtubule capture and attachment at the kinetochore (Abrieu et al. 2000; Yao et al. 2000). Unsurprisingly, *Cenpe* deletion in mice is early embryonic lethal (Putkey et al. 2002). Conditional *Cenpe* deletion in mouse embryonic fibroblasts and an adult regenerating liver system resulted in cells with profound mitotic defects including chromosome misalignment and segregation failure (Putkey et al. 2002; Weaver et al. 2003). Firstly, we detail the clinical presentation and progression of both siblings. Then, using patient-derived lymphoblastoid cell lines (LCLs), we catalog a series of mitotic abnormalities including aberrant spindle microtubule organization, delayed mitotic progression and elevated levels of binucleate cells. The latter phenotype, in particular, is suggestive of an impaired ability to exit mitosis effectively. We also find that the nuclei of these binucleates are often of unequal size indicative of impaired chromosome segregation during mitosis and subsequent cytokinesis failure. To further characterize and consolidate the cellular pathogenicity of the *CENPE* variants we have identified, we model each individually and in combination using Flip-In technology (Invitrogen) coupled with siRNA-mediated knockdown of the endogenous CENP-E in an independent cell system.

Interestingly, we find that many of these cellular phenotypes observed in the patient LCLs are also observed in LCLs from a patient with a pathogenic defect in PCNT; the centrosome-associated protein implicated in MOPDii (Griffith et al. 2008; Rauch et al. 2008). Therefore, using MPD patient-derived material, we show that a novel kinetochore-associated defect in CENP-E shares overlapping phenotypes of abnormal spindle microtubule structure and mitotic progression to that of a pathogenic defect in a centrosome-associated protein.

Methods

DNA extraction

Genomic DNA was extracted from peripheral blood samples using either the Puregene kit[®], Magnapure[®] or Autogen[®] systems following the manufacturers' recommendations.

Whole exome sequencing and analysis

We performed whole exome sequencing (WES) of peripheral blood DNA from subject LR05-054a1. We used the Nimblegen whole exome capture kit, and sequence was generated on an Illumina GAI machine. Sequence was aligned to hg19 using BWA 0.6.2 and single nucleotide variants and indels were called using GATK 2.3.9 UnifiedGenotyper. Mean coverage was calculated using GATK 2.3.9 Depth Of Coverage Walker. Annotation of variants, including identification of variants present in dbSNP, 1,000 Genomes, and the NHLBI Exome Variant Server (EVS), was performed with Annovar (Wang et al. 2010). Pedigree-specific nonsynonymous variants, including potential compound heterozygotes, were identified using SOLVE-Brain. Variants in genes associated with known PM and Seckel-related syndromes, as well as other known developmental brain disorders, were identified using SOLVE to filter against the contents of the Developmental Brain Disorders Database 1.2 (DBDB). Final annotation of biological function and pathway membership of candidate genes with nonsynonymous compound heterozygous variants were performed using Lynx (Sulakhe et al. 2014).

Weblinks:

BWA: <http://bio-bwa.sourceforge.net/bwa.shtml>.

GATK: http://www.broadinstitute.org/gsa/wiki/index.php/Home_Page.

Home_Page.

Annovar: <http://www.openbioinformatics.org/annovar/>.

SOLVE: <https://github.com/alex-paciorkowski/SOLVE>.

DBDB: <https://www.dbdb.urmc.rochester.edu/associations/list>.

Lynx: <http://lynx.ci.uchicago.edu:8080/>.

Sanger sequencing

PCR amplification was performed with 50 ng of genomic DNA using Taq DNA polymerase (Applied Biosystems[®]). Exons 23 and 29 of *CENPE* (NM_001813.2) were targeted for mutation analysis by Sanger sequencing using standard methods.

Cell lines

EBV transformed lymphoblastoid cell lines (LCLs) were made from both patients (LR05-054a1 and LR05-054a2)

using lymphocytes fractionated from whole blood. Wild-type (WT; AG09387) cells were obtained from Coriell Cell Repository (New Jersey, USA). *PCNT*-mutated MOPDii LCLs were from patient CV1559 and carry a homozygous mutation in *PCNT* resulting in p.E220X; Family 1 from Griffith et al. (2008).

Antibodies

Anti-53BP1 was the mouse monoclonal from Bethyl. Anti-CENP-E was mouse monoclonal [1H12] from Abcam used for patient LCL indirect immunofluorescence whilst anti-CENP-E (AKIN04) from cytoskeleton was used for Western blotting. Anti-BUBR1 was a mouse monoclonal (ab4637) from Abcam. Anti- α -tubulin was a mouse monoclonal (clone B512) from Sigma-Aldrich and anti- β -tubulin was a rabbit polyclonal (H-235) from Santa Cruz. Anti-PARP (MCS 1522G) was from AbD Serotec and anti-p85-PARP (cleaved-PARP: G7341) was obtained from Promega.

Indirect immunofluorescence

Following various treatments (see below), LCLs were pelleted, swollen in 75 mM KCL (10 min), fixed in 4 % paraformaldehyde (10 min) prior to cytospinning onto poly-L-lysine coated slides. LCLs were then permeabilized (0.1 % Triton X-100 in 5 % BSA-PBS for 2 min) and blocked (5 % BSA-PBS for 10 min) prior to incubation with primary and secondary antibodies diluted in 5 % BSA-PBS. Slides were counter-stained with 4',6-diamidino-2-phenylindole (DAPI) and mounted in Vectashield anti-fade medium. Images were captured using SimplePCI software on the Zeiss Axioplan platform.

LCL treatments

HU-induced 53BP1 foci

LCLs were treated with 5 mM HU for 2 h prior to harvesting and processing for indirect immunofluorescence staining using anti-53BP1.

Microtubule spindle stabilization

Spindles were stabilized by treating exponentially growing LCLs with Taxol (10 μ M) and the proteasome inhibitor MG132 (10 μ M) for 1 h in complete medium at 37 °C before shifting the cells to packed ice for 1.5 h. LCLs were then pelleted and processed for indirect immunofluorescence using anti- α -tubulin.

BubR1 phosphorylation

Exponentially growing LCLs were treated with 0.2 µg/ml colcemid for 2 h prior to pelleting and whole cell extracts made using urea buffer (9 M urea, 50 mM Tris-HCl at pH 7.5 and 10 mM 2-mercaptoethanol) and sonication (15 s at 30 % amplitude using a micro-tip; Sigma-Aldrich).

Colcemid block and release into cytochalasin B

LCLs were treated with colcemid (0.2 µg/ml for 4 h), pelleted, washed in complete medium before release into complete medium supplemented with cytochalasin B (5 µg/ml). Cells were incubated for 24 h and processed for indirect immunofluorescence using anti-β-tubulin.

G2-M checkpoint analysis

LCLs were irradiated with 3 Gy IR in complete medium or 7 J/m² in PBS before seeding into complete medium supplemented with 0.2 µg/ml colcemid and incubated for 4 h. LCLs were pelleted, swollen in 75 mM KCL (10 min) and fixed in 4 % paraformaldehyde (10 min) prior to cytospinning onto poly-L-lysine coated slides. Cells were counterstained with DAPI and mounted in Vectashield anti-fade prior to analysis on the Zeiss Axioplan system.

Flp-In system and variant modeling

The Flp-In (Invitrogen) system employed here for *CENPE* in DLD-1 cells has been described previously (Kim et al. 2010). The cDNA and siRNA for *CENPE* used here are identical to those of Kim et al. (2010). Lipofectamine RNAi MAX (Invitrogen) was used for *CENPE* siRNA (5'-CCACUAGAGUUGAAAGAU-3') with doxycycline used at 1 µg/ml for induction of siRNA-resistant *CENPE* construct. For the live cell imaging, an RFP-histone H2B expressing DLD-1 line was used to visualize chromosomes, as described in Kim et al. (2010).

Results

Clinical presentation and progression of siblings LR05-054a1 and LR05-054a2

The features of both siblings are summarized in Table 1. This family is a non-consanguineous family of European ancestry. The first affected, a boy (LR05-054a1), was born at 34 weeks of gestation to 33-year-old mother, who had a history of one miscarriage. Pregnancy was complicated by intrauterine growth restriction detected at 6 and 10 weeks of gestation; biparietal diameter, head circumference,

femoral length, abdominal circumference and estimated fetal weight were all less than 10th percentile for gestational age. At 34 weeks of gestation, prenatal ultrasound was performed due to maternal hypertension and showed a severely small fetus for gestational age with decreased fetal activity, so delivery was induced at that time. At birth, LR05-054a1 exhibited severe congenital microcephaly, with a birth OFC of 24.5 cm (−5 SD), growth deficiency with a weight of 1.5 kg (−3 to −4 SD) and length 40.5 cm (−2 to −3 SD). Facial features included a sloping forehead, prominent nose, and mild micrognathia. This boy had a single seizure at 7 months of age which consisted of generalized body stiffening, extension of the arms and legs with breath holding. Electroencephalogram was abnormal showing generalized rhythmic fast activity followed by generalized slowing, and multifocal sharp waves. He had a few additional seizures between 7 and 10 months of age that were subsequently well controlled on phenobarbital.

At 4 months of age, he had a viral upper respiratory illness and, because of worsening respiratory symptoms, was admitted and underwent a cardiac echocardiogram which showed abnormal tricuspid and mitral valve inflow patterns with increased early and late atrial (e/a) ratios, suggestive of diastolic dysfunction, and decreased deceleration times varying with the respiratory cycles, indicative of increased flow reversal in the pulmonary veins with atrial contraction. Additional echo findings include trivial tricuspid and mitral regurgitation, biatrial enlargement with paradoxical septal wall motion suggesting elevated right ventricular systolic pressure, a trabeculated and hypertrophied apex of the left ventricle but with normal dimensions and good systolic function. Overall, these features were consistent with congenital restrictive cardiomyopathy, for which he was treated with digoxin, Lasix and aspirin. Other medical complications include poor weight gain, partly due to the underlying syndrome and partly due to gastroesophageal reflux, hyperopia requiring glasses, and mild congenital subglottic stenosis of no clinical significance.

On last examination at 5 years, LR05-054a1 exhibited the following growth measurements: weight 9.14 kg (−5 SD), length 83.7 cm (−7 SD), and OFC 37.0 cm (−9 SD). His facial appearance was significant for striking microcephaly with a very low forehead, round face, prominent nose, mild micrognathia and decreased subcutaneous fat (Fig. 1a–c). His head appeared disproportionately smaller than his body whilst his nose and ears disproportionately large. His hands and feet were mildly small. Even though LR05-054a1 did not exhibit the nasal preponderance with downward curved nasal tip classically associated with Seckel syndrome, his overall features appeared compatible with profound MPD of this type (Hall et al. 2004; Majewski et al. 1982a, b). On neurological exam, he was alert but did not follow commands or communicate. Eye movements

Table 1 A summary of the clinical and neuroimaging features of the two siblings

Features	LR05-054a1	LR05-054a2
Gender	Male	Female
Age last assessed	8 years	3 years
Growth at birth		
OFC (SD)	−5	nd
Weight (SD)	−3 to −4	−3
Length (SD)	−2 to −3	−3
Postnatal-age	5 years	3 years
OFC (SD)	−9	−7
Weight (SD)	−5	−3 to −4
Length (SD)	−7	−2 to −3
Facial features	Apparent MIC, prominent nose, sloping forehead, relatively large ears, mild micrognathia	Apparent MIC, low-sloping forehead, round face, proportionately large ears and nose
Extremities	Mildly small hands and feet	Small hands and feet (−2 SD below mean)
Cardiac abnormalities	Congenital restrictive cardiomyopathy	None
Seizures	Tonic seizures, onset 7 months	None
Tone	Mild lower extremity spasticity	Mildly increased
Development	Severe DEV delay, nonverbal	Less severe DEV delay, 65 % language delay at 1 year 4 months
Other medical problems	GER, mild congenital subglottic stenosis	Otitis media, tight heel cords
Neuroimaging features	Simplified gyral pattern, thin cortex, mildly disproportionate cerebellar hypoplasia	nd
Outcome	Deceased at 8 years (pneumonia)	Alive

DEV developmental, GER gastroesophageal reflux, MIC microcephaly, nd no data, OFC occipitofrontal circumference, SD standard deviation

were full, pupils normal and facial movements symmetric. Motor exam showed mild spasticity in the hip adductors and ankles. Deep tendon reflexes were normal (2+) and symmetric. Developmentally, this child made very slow but steady progress albeit remaining severely handicapped. He began to pull himself to stand only briefly at 3.5 years of age, and crawled at 4 years. On assessment at 5 years, he had no communication skills besides vocalizing without signs, words or pointing. He used a rake and pincer grasp. He was fed a soft diet by mouth. His general health including his cardiac course deteriorated over time; he developed pericardial effusion on a few occasions requiring percutaneous drainage and also had several episodes of pneumonia, with progressive worsening of his oxygen saturation levels. A final episode of pneumonia at 8 years of age proved fatal.

The pregnancy of the second affected child of the family, a girl (LR05-054a2), was also complicated by microcephaly and intrauterine growth restriction prenatally detected on ultrasound. She was born at 37 weeks of gestation with birth weight 1.76 kg (−3 SD) and length 41.5 cm (−3 SD). She was noticed to have the same facial features as her brother's at birth. She exhibited frequent episodes of otitis media requiring myringotomy tube placement, and minor foot deformities consisting of tight heel cords

and medially deviated halluces, for which she underwent heel cord lengthening and tenotomy of great toes for better alignment. Unlike her brother, she did not have episodes of pneumonia or other severe illnesses requiring hospitalization. Echocardiogram and electrocardiogram were normal. She did not have seizures, and was described as a generally happy child.

On last exam at 3 years, her weight was 9.1 kg (−3 to −4 SD), length 83 cm (−2 to −3 SD), and OFC 37.5 cm (−7 SD). Her palms, fingers and toes measured less than −2 SD. Her overall appearance was significant for a small head, small round face with a low forehead and proportionately large ears and nose (Fig. 1d). The helix of the right ear was mildly under-folded, and great toes medially deviated. Neurological exam revealed an alert child with brief visual tracking and a strong cry. Tone was mildly increased and deep tendon reflexes minimally brisk (2++). Eye movements were full with intermittent nystagmus. Facial movements were symmetrical. Developmentally, this girl sat alone at 1–1.5 years, pulled to stand, cruised and climbed on furniture at 2–2.5 years, and crawled at 2.5 years. At 2.5 years, her language was limited to a few words such as “hi” and “bye”, along with mimicking speech sounds. She picked small objects but did not point or draw. She attended preschool at 3 years of age. She was able to chew solids

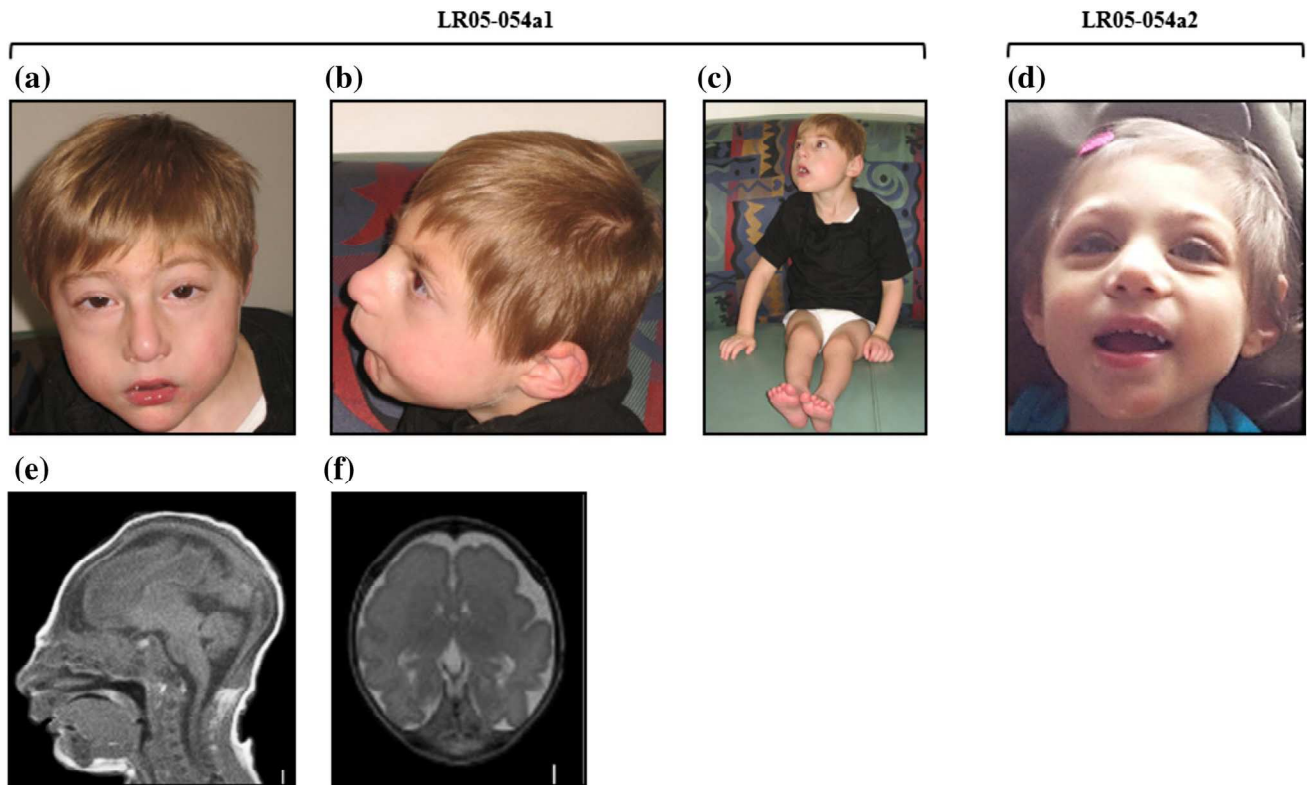


Fig. 1 Images of the affected siblings LR05-054a1 and LR05-054a2. **a–c** Patient LR05-054a1 at 16 months of age. **d** Patient LR05-054a2 at 2 years of age showing microcephaly, low-sloping forehead, proportionately large ears and nose, a prominent nose, micrognathia, and severe growth deficiency. **e** T1-weighted mid-sagittal and **f**

T2-weighted axial brain MRI images of LR05-054a1 at 17 days of life show severe microcephaly, with low-sloping forehead, severely simplified gyral pattern with virtually no gyri in the frontal region, mildly increased extra-axial space, and disproportionately small cerebellar vermis

with some feeding difficulties, such as choking occasionally, requiring a softer diet.

Clinical work-up of siblings LR05-054a1 and LR05-054a2

Clinical workup performed on LR05-054a1 with inconclusive results includes a karyotype, a chromosomal microarray (Signature Genomics), chromosome breakage studies, metabolic tests and TORCH titers. A skeletal survey demonstrated apparent microcephaly with no evidence of craniosynostosis, subtle widening of the ribs and possible metaphyseal areas of sclerosis in the distal femurs and proximal tibias. The hands appeared osteopenic and metacarpals were relatively short. Overall, these findings were not characteristic of a specific bone dysplasia. Brain MRI scan of LR05-054a1 at 17 days of life showed extreme microcephaly with a low forehead and a diffuse severely simplified gyral pattern with virtually no gyri over the frontal lobe, a thin 2–3 mm cortex and immature white matter (Fig. 1e, f). Additional findings include partial agenesis of the corpus callosum and cerebellar hypoplasia that appeared

disproportionate to the degree of microcephaly (Fig. 1e, f). Notably, neither sibling had a history of blood dyscrasias or other childhood cancers.

Molecular-genetic analysis of family LR05-054

Although LR05-054a1 did not present with the skeletal features characteristic of MOPDii (e.g., coxa vara, small iliac wings, V-shaped distal femoral epiphysis, metaphyseal flaring of radius and ulna), these features can take time to manifest (Hall et al. 2004). Since defects in the centrosome-associated protein pericentrin, encoded by *PCNT*, are currently one of the most commonly identified defects associated with profound MPD in humans (MOPDii) (Bober et al. 2012), we sequenced *PCNT* and did not find any pathogenic mutations in this family.

WES was then performed using peripheral blood from subject LR05-054a1. Mean depth of coverage across the targeted exome was 99.5x. The mean depth of coverage across *CENPE* was 45.5x. SOLVE identified 4,322 genes with two or more heterozygous nonsynonymous variants.

However, most of these variants were in genes not likely contributory to the phenotype (DBDB, dbSNP and EVS). There were no novel variants detected in other known PM, MPD or Seckel syndrome genes.

Annotation of biological function and pathway membership in Lynx allowed us to identify compound heterozygous nonsynonymous variants in *CENPE* (c.2797G>A;p.D933N and c.4063A>G;p.K1355E) as putatively causative of the phenotype (Fig. 2a). Although both of these variants were present in dbSNP and the EVS, it was with very low frequency (1/8597 for c.2797G>A;p.D933N and 3/8595 for c.4063A>G;p.K1355E), not ruling out their involvement in an autosomal recessive developmental disorder. Two additional *CENPE* variants (chr4: 104059542 encoding c.6269C>T;p.T2090M and chr4: 104066461 encoding c.4603T>C;p.F1535L) were identified; however, as these were both found in high frequency in dbSNP and the EVS they were not considered to be pathogenic. We, therefore, proceeded with confirmatory studies for the *CENPE* c.2797G>A;p.D933N and c.4063A>G;p.K1355E variants. Subsequent Sanger sequencing confirmed that the parents (LR05-054F: father; LR05-054M: mother) were found to each carry one of these mutations (Fig. 2a). Mutation status of both siblings was also confirmed by Sanger sequencing (Fig. 2a). Both residues (D933 and K1355) are relatively highly conserved in mammals, although not in the mouse *M. musculus* (Fig. 2b). Patient-derived LCLs were prepared from both siblings for functional analysis. Sanger sequencing of *CENPE* genomic DNA from ten clinically overlapping individuals did not reveal any mutations (See Online Resource Table 1 for a summary).

LR05-054a1 mitotic LCLs exhibit reduced centromeric accumulation of CENP-E

CENP-E is required for spindle microtubule capture to the kinetochore (Fig. 2c). Its expression typically peaks during G2-M and is then rapidly degraded in telophase (Brown et al. 1994). CENP-E expression by Western blotting was comparable in whole cell extracts from asynchronous wild-type (WT) LCLs to those LCLs from both siblings (LR05-054a1 and LR05-054a2) and *PCNT*-mutated MOPDii, suggesting that the variants did not adversely impact on CENP-E expression or stability (Fig. 3a). As expected, the majority of WT mitotic cells exhibited a strong bivalent CENP-E signal at chromosome centromeres (Fig. 3b–d). This was in stark contrast to LCLs from LR05-054a1 where approximately 80 % of mitotic cells exhibited a barely detectable CENP-E signal (Fig. 3b–d). Interestingly, *PCNT*-defective MOPDii LCLs also exhibited a reduced intensity of bivalent CENP-E signal in about 50 % of mitotic cells, highlighting a

phenotypic similarity between CENP-E mutant cells and those of MOPDii (Fig. 3d).

LR05-054a2 LCLs exhibit impaired BubR1 phosphorylation

Phosphorylation of the kinetochore-associated kinase BubR1 occurs at unattached kinetochores as part of the SAC. This has been proposed to represent autophosphorylation that is dependent upon CENP-E (Guo et al. 2012; Mao et al. 2005; Tanudji et al. 2004). Although others have argued that phosphorylation must be mediated by other kinases as BubR1 itself has been proposed to be a pseudokinase without intrinsic kinase activity (Suijkerbuijk et al. 2012). Regardless, mitotic-BubR1 phosphorylation can be monitored as an overt electrophoretic retardation following treatment with spindle poisons (Tanudji et al. 2004). As an indirect assay of CENP-E functional proficiency, we treated LCLs with colcemid for 2 h and examined the electrophoretic mobility of BubR1 by Western blotting. Colcemid treatment of WT LCLs resulted in the expected retardation in BubR1 mobility, in contrast to LR05-054a2 LCLs (Fig. 3e). Interestingly, *PCNT*-MOPDii LCLs, similar to LR05-054a2, failed to induce a shift in BubR1 mobility suggestive of impaired CENPE-dependent BubR1 phosphorylation under these conditions (Fig. 3e). These results are consistent with the reduced CENP-E localization to mitotic centromeres observed in both these LCLs (Fig. 3d).

LR05-054a1 LCLs exhibited elevated sensitivity to Taxol

Prolonged treatment of LR05-054a1 LCLs for up to 24 h with the spindle poison Taxol was associated with elevated levels of apoptosis compared to WT LCLs, as revealed by elevated p85-PARP expression levels. These data demonstrate that these patient cells are sensitive to treatment with a microtubule spindle stabilizing agent.

LR05-054a1 LCLs exhibit altered spindle dynamics

Through its role in chromosome congression and spindle capture at the kinetochore, CENP-E enables chromosomal alignment prior to effective segregation (Fig. 2c). Occasionally, lagging chromosomes were evident in mitotic cells from untreated asynchronous LR05-054a1 cells, but the frequency of this was quite variable (Fig. 4a). Nevertheless, these were not seen in over 100 mitotic cells analyzed from WT LCLs. We next examined the structure of the microtubule spindle network following stabilization with a short pulse of Taxol via indirect immunofluorescence with antibodies against α -tubulin. Most (>80 %) of the WT mitotic cells exhibited a bipolar spindle (Fig. 4b,

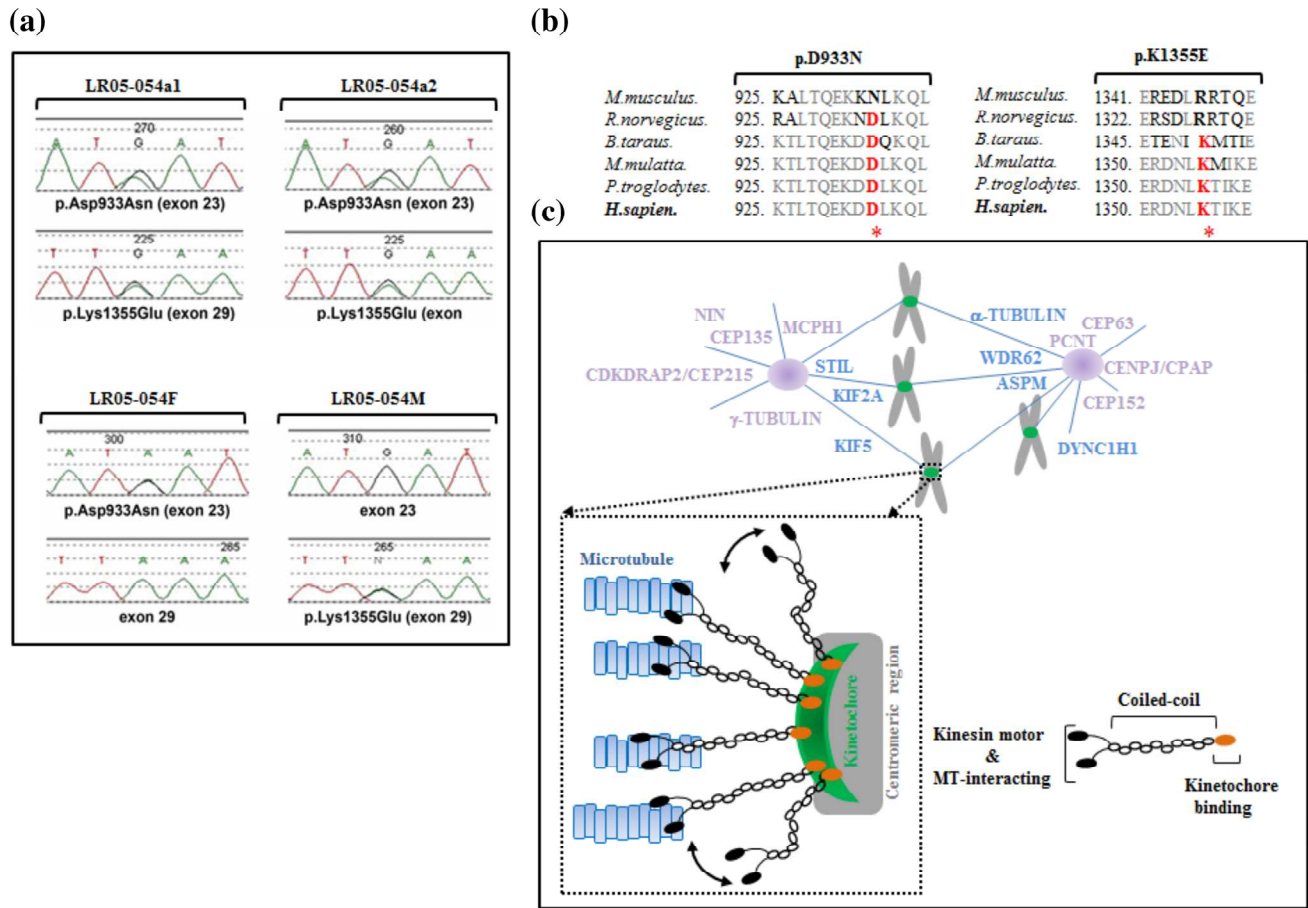


Fig. 2 Family LR05-054 have compound heterozygous variants in *CENPE*. **a** Representative sequence chromatographs identifying mutations in *CENPE* (NM001813.2) showing the two missense mutations identified in both siblings (c.2797G>A, p.D933N and c.4063A>G, p.K1355E; a-b), as well as parental carrier status: p.D933N is paternally inherited (LR05-054F; father) and p.K1355E is maternally inherited (LR05-054M; mother). **b** Multiple sequence alignments of the regions around amino acids 933 and 1,355 were assembled using MUSCLE (version 3.6) via NCBI. Identical sequences are identified in gray, divergent in black and the target residues in red and by the red asterisk. Both D933 and K1355 are relatively well conserved in mammals. Although interestingly, neither residue is conserved in mice, *M. musculus*. In fact, the mouse contains N933, one of the variants identified in our patients. It is notable that CENP-E protein identity between *H. sapien* and Cenpe of *M. musculus* is only 64.4 % (for *R. noregicus* 56 %). This is in marked contrast to chimpanzee *P. troglodytes* (98.8 %), Rhesus monkey *M. mulatta* (95.8 %) or cattle *B. taurus* (79 %). In fact, the mouse (*M. musculus*) Cenpe is only 2,471 amino acids in length compared to

2,701 amino acids of the human CENP-E. Whilst both proteins possess the kinesin motor domain ATPase, only the human protein possesses the SMC (C0G1196) chromosome segregation ATPase and SMC_N(pFam02463) RecF/RecN/SMC N-terminal domain, highlighting the significant sequence and domain disparity between mouse and human herein. **c** Schematic representation of a mitotic cell, showing the microtubule nucleating centrosome (purple), the microtubule spindles (blue), mitotic chromosomes (gray) and their centromeric kinetochores (green). Known genetic defects in centrosomal and centrosome-associated proteins are shown in purple and spindle pole components in blue. The dashed box indicates a magnified schematic of the kinetochore showing CENP-E capturing microtubules. CENP-E is characterized by a large 230-nm flexible coiled-coil domain that can move about to enable microtubule capture by the globular head regions which are composed of microtubule interaction, kinesin motor and ATP-binding domains. The *CENPE* mutations identified here encode residues within this coiled-coil (color figure online)

c). In direct contrast, approximately 80 % of mitotic cells from PCNT-MOPDii (PCNT) and LR05-054a1 exhibited multipolar spindles suggestive of massively disorganized spindle network in these cells (Fig. 4b, c). Abnormal spindle structure has been reported for PCNT-MOPDii previously (Rauch et al. 2008). Interestingly, again we observe similar cellular phenotypes between a centrosomal defect (PCNT) and that of *CENPE*-mutated LR05-054a1 LCLs.

LR05-054a2 LCLs exhibit elevated levels of binucleates with features consistent with altered chromosome segregation

CENP-E deficiency is associated with elevated frequency of polar chromosomes and mis-segregation (Guo et al. 2012; Putkey et al. 2002; Tanudji et al. 2004; Weaver et al. 2003). We did not observe a significantly elevated

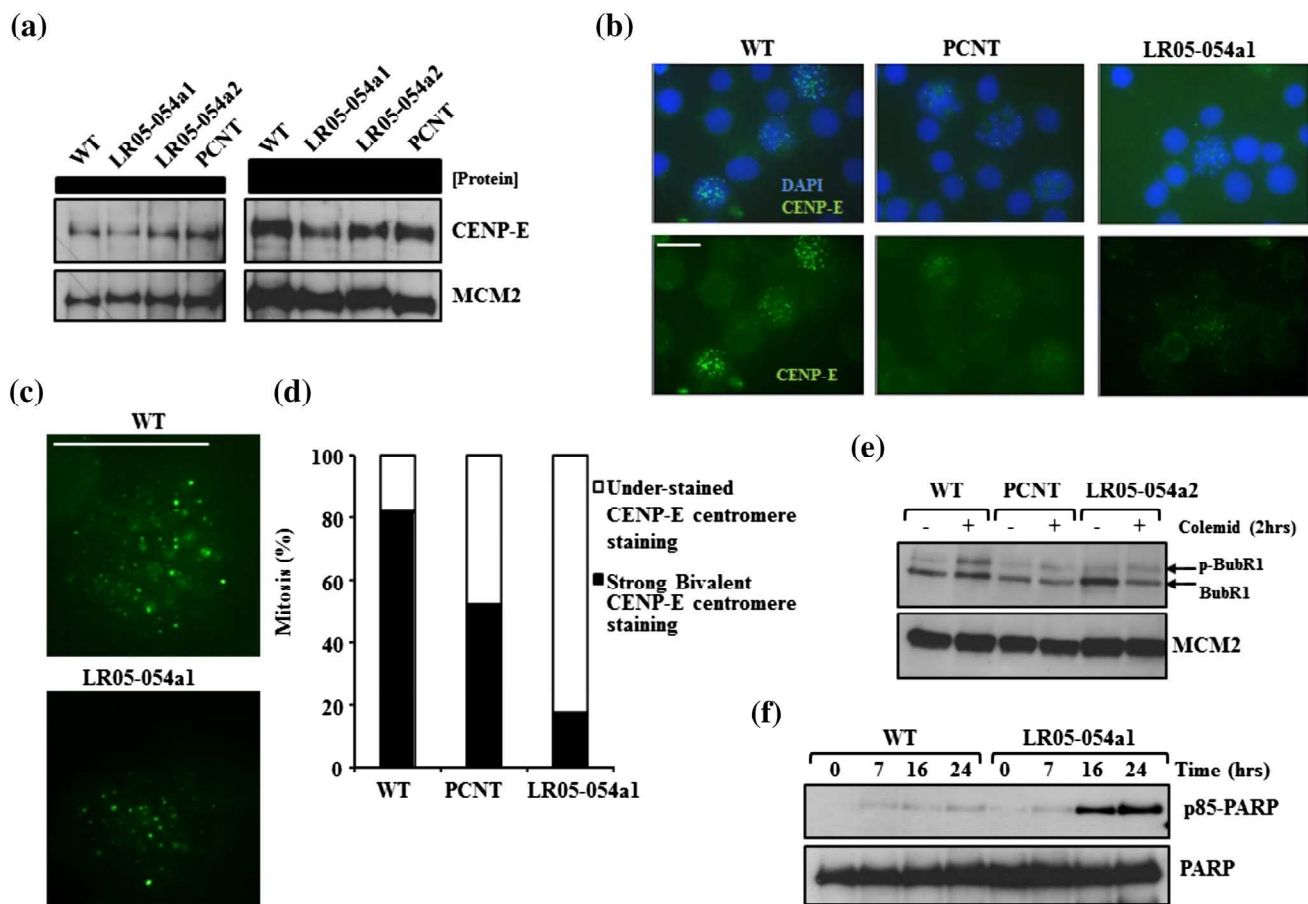


Fig. 3 CENPE- and PCNT-mutated patient LCLs exhibit reduced kinetochore localization of CENP-E and reduced BubR1 phosphorylation. **a** CENP-E expression is comparable in whole cell extracts (WCE) from wild-type (LCLs) and LCLs from both siblings (LR05-054a1 and LR05-054a2) and from PCNT-mutated MOPDii LCLs. The left-hand panels show 25 μ g WCE and the right-hand panel 50 μ g of WCE. **b** Indirect immunofluorescence (IF) showing CENP-E focalization onto kinetochores of mitotic cells in wild-type (WT), PCNT-mutated MOPDii (PCNT) and CENPE-mutated patient derived LCLs (LR05-054a1). Cells were treated with Taxol (10 μ M) and MG132 (10 μ M) for 1 h followed by 1.5 h on ice to stabilize spindle microtubules. Both PCNT and LR05-054a1 mitotic cells exhibit a clear under-representation of defined CENP-E foci. All images were acquired under identical exposure conditions (scale bar; 10 μ m). **c** Close-up image of CENP-E focalization in a mitotic cell from wild-type (WT) and CENPE-mutated patient derived LCLs (LR05-054a1). It has been estimated that only approximately 50 CENP-E dimers occupy each normal human kinetochore (Brown et al. 1994) (scale bar; 10 μ m). **d** The distribution of mitotic cells exhibiting strong bivalent CENP-E foci and those under-stained for

CENP-E foci in wild-type (WT), PCNT-mutated MOPDii (PCNT) and CENPE-mutated patient derived LCLs (LR05-054a1). Between 30 and 50 mitotic cells were scored for each cell line under identical conditions as **a**. Strong bivalent CENP-E staining is under-represented in PCNT and LR05-054a1 LCLs compared to WT. **e** LCLs, from wild-type (WT), PCNT-mutated MOPDii (PCNT) and the female CENPE-mutated patient sib (LR05-054a2) were treated with colcemid (0.2 μ g/ml) for 2 h and phosphorylation status of BubR1 was determined by electrophoretic shift following Western blotting. In contrast to WT, colcemid treatment did not induce significant BubR1 retardation (phosphorylation) in PCNT or LR05-054a2 LCLs under these conditions. Interestingly, following a more prolonged treatment with colcemid (≥ 4 h), BubR1 retardation is evident in both these LCLs suggestive of a leaky/hypomorphic defect. The blot was re-probed for MCM2 expression as a loading control. **f** LCLs were treated with 2.5 μ M Taxol and incubated for the times indicated. Apoptosis induction and levels were assessed by blotting for cleaved form of PARP; p85-PARP. The p85-PARP was detected at 16 h post-treatment in LR05-054a1 LCLs and further increasing at 24 h. Under these conditions, no apoptosis was evident in the WT LCLs

spontaneous frequency of micronuclei in PCNT or CENPE LCLs (LR05-054a2), although we did detect an approximately two-fold increase in the level of bi-nucleated cells here, compared to WT (Fig. 4d). Binucleates are the product of impaired and/or delayed cytokinesis which can be a consequence of a failure to coordinately transit mitosis

effectively. Strikingly, approximately 50 % of binucleates detected in the LR05-054a2 LCLs exhibited nuclei of unequal size, suggestive of unequal segregation during mitosis and impaired cytokinesis (Fig. 4e). We did not observe this phenotype in the binucleates from the PCNT-MOPDii LCLs.

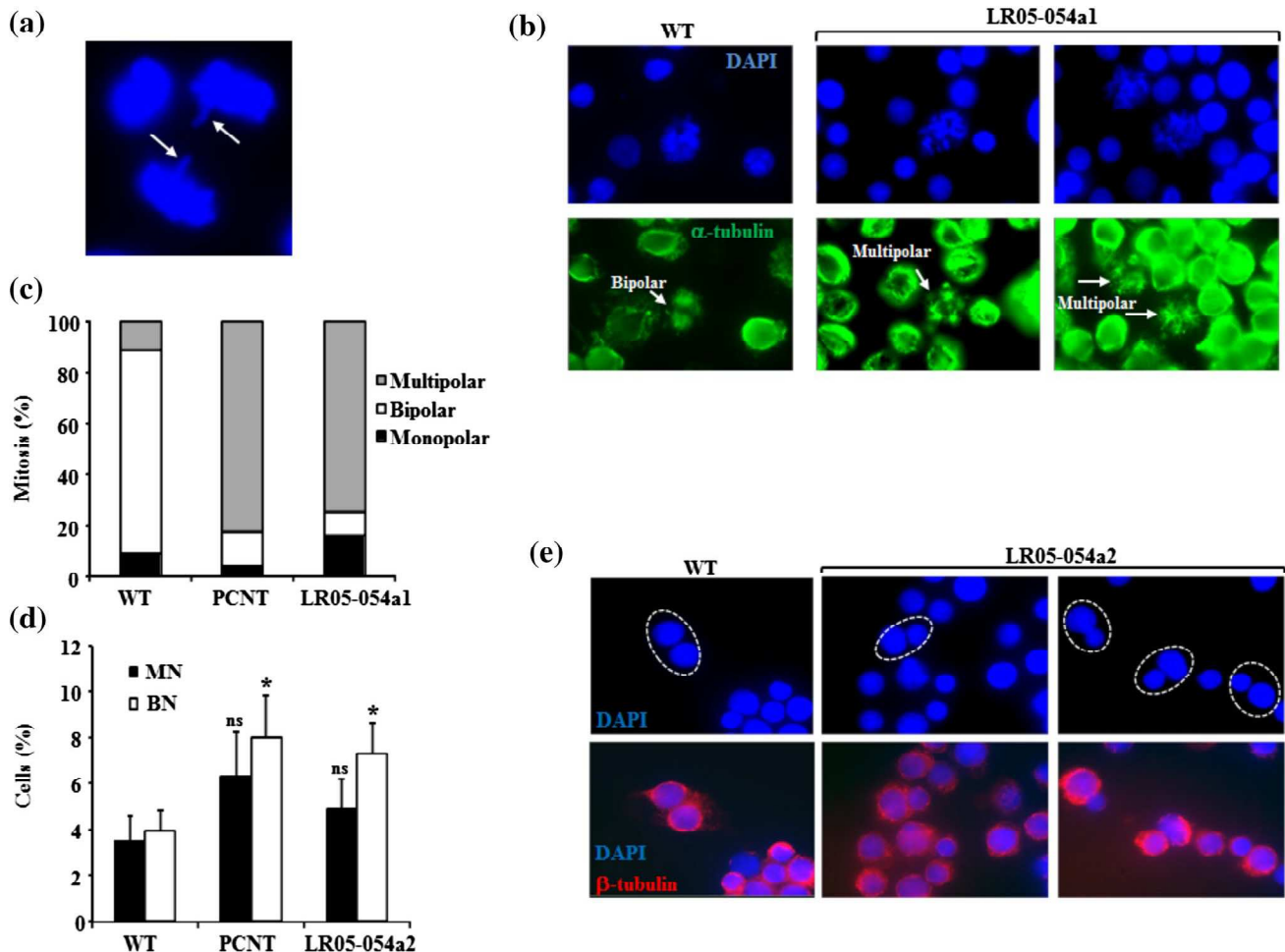


Fig. 4 CENPE-mutated patient LCLs exhibit abnormal spindle microtubule organization and evidence of aberrant segregation and delayed progression through mitosis. **a** An image of a lagging chromosome in an anaphase cells from an asynchronous culture of LR05-054a1. **b** The lower panels show IF images of α -tubulin staining microtubule network (green) from wild-type (WT) and LR05-054a1 LCLs following treatment with Taxol (10 μ M) and MG132 (10 μ M) for 1 h then 1.5 h on ice to stabilize spindle microtubules. The upper panels are the corresponding DAPI-staining nuclei. A normal bipolar spindle is seen in WT mitotic cell whilst multipolar spindles are frequently seen in mitotics from LR05-054a1. **c** The distribution of spindle abnormalities noted in mitotic cells in wild-type (WT), PCNT-mutated MOPDii (PCNT) and CENPE-mutated patient derived LCLs (LR05-054a1). Between 40 and 50 mitotic cells were scored for each

cell line. Multipolar spindles are over-represented in mitotic cells from PCNT and LR05-054a1 LCLs compared to WT. **d** The spontaneous levels of micronuclei (MN) are not statistically significantly elevated (ns not significant Student *t* test) in PCNT-mutated MOPDii (PCNT) or CENPE-mutated patient derived LCLs (LR05-054a2) compared to WT. In contrast, binucleates (BN) levels are approximately twofold higher in these LCLs compared to WT potentially suggestive of elevated rates of cytokinesis failure (**p* < 0.05 Student *t* test). **e** DAPI (upper panel) and their corresponding IF-mediated β -tubulin images (lower panel) of binucleates (circled) from LR05-054a2 LCLs frequently exhibit nuclei of different sizes. The β -tubulin staining facilitates the identification of true binucleate cells. Differing sized nuclei within a binucleate are indicative of unequal segregation during mitosis (color figure online)

Expression of the D933N and K1355E variants of CENP-E induce mitotic spindle abnormalities and polar chromosomes

We used Flp-In (Invitrogen) recombination to introduce siRNA-resistant p.D933N and p.K1355E variants of CENPE, individually and in-cis (Double mut; double mutant), into DLD-1 cells under doxycycline (Dox) conditional regulation, as described in detail by Kim et al. (2010). Following siRNA-mediated silencing of the

endogenous CENPE, Dox treatment (8 h) resulted in strong expression of these variants and cells were fixed for immunofluorescence imaging. This allowed us to interrogate the impacts on mitotic spindle structure and chromosome segregation when each variant was expressed individually and when engineered together into CENP-E. The induction scheme used is shown in Fig. 5a and comparable CENP-E expression levels shown in Fig. 5b. In this independent model cell system, we found that p.D933N and p.K1355E individually or when in-cis caused an elevated frequency

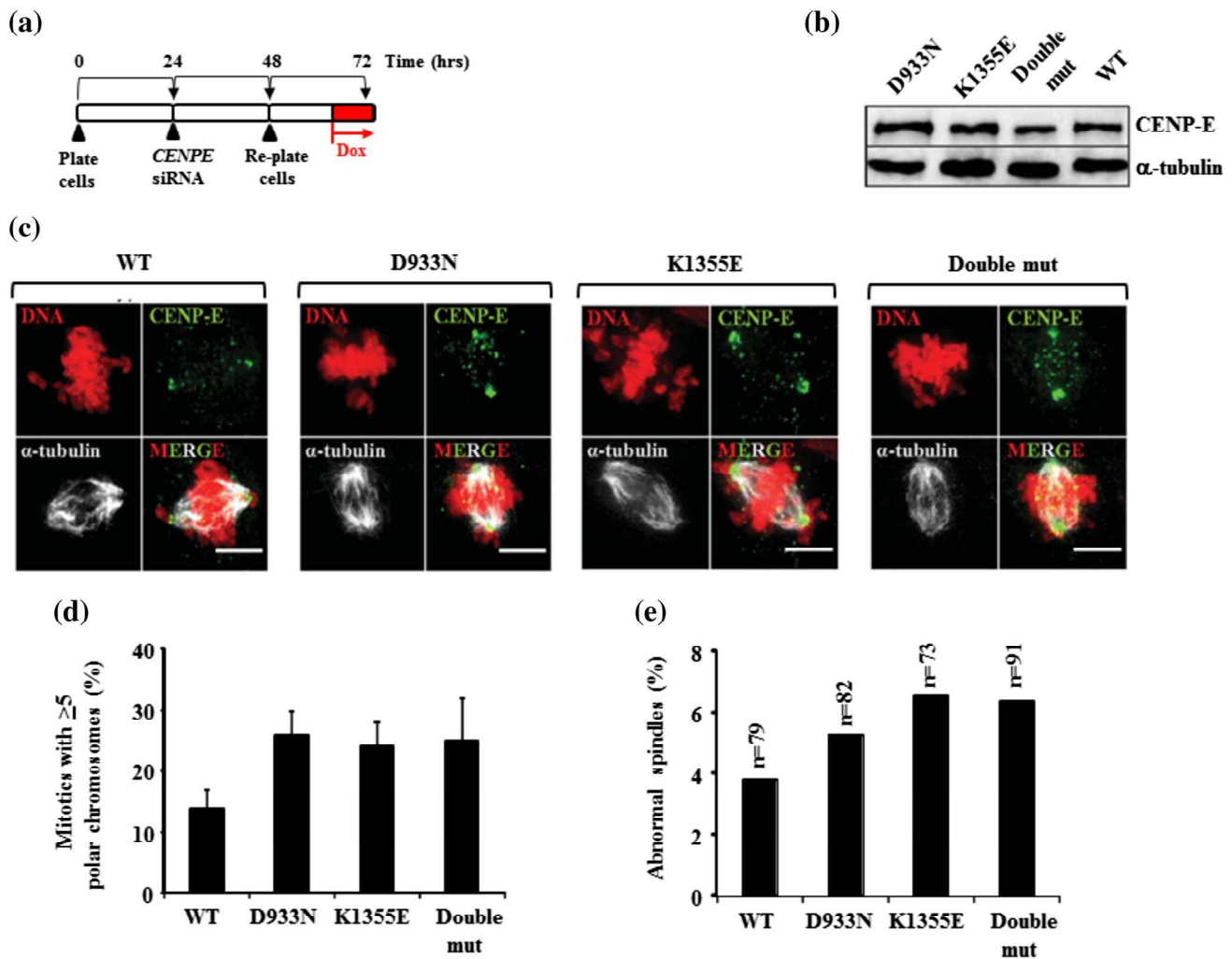


Fig. 5 Expression of the D933N and K1355E variants of CENP-E results in mitotic spindle abnormalities and polar chromosomes. **a** DLD-1 cells engineered by Flp-In recombination for the conditional (doxycycline-induced) expression of siRNA-resistant *CENPE* and *CENPE* patient variants were plated and 24 h later subjected to *CENPE* siRNA targeting endogenous CENP-E only. After 40 h approx., the cells were treated with doxycycline (Dox; 1 μ g/ml) for 8 h to induce the expression of the siRNA-resistant *CENPE* variants. Cells were then fixed and processed for immunofluorescence. **b** Expression levels of the various CENP-E variants including a construct with both variants engineered in-cis (Double mut; double mutant) were compared to that of wild-type (WT) CENP-E following 8 h in doxycycline. **c** Immunofluorescence images following 8 h treatment with doxycycline for wild-type (WT), D933N alone, K1355E

alone and the double mutant (Double mut) with both D933N and K1355E in-cis co-stained with anti-CENP-E and anti- α -tubulin. Polar chromosomes not aligned properly stain strongly for CENP-E under these conditions as seen in D933N alone, K1355E alone and the double mutant (Scale bar 5 μ m). **d** The frequency of mitotic cells with ≥ 5 unaligned polar chromosomes is elevated following the expression of D933N alone, K1355E alone or following the expression of both variants in-cis (Double mut). Between 70 and 90 mitotic cells were evaluated in total for each construct. **e** Expression of D933N alone, K1355E alone or both variants in-cis (Double mut) resulted in an increased frequency of mitotic cells with abnormal spindle structure (monopolar plus multipolar spindles) compared to the expression of wild-type (WT) CENP-E in this system (*n* the number of mitotic cells scored)

of mitotic cells with polar chromosomes (Fig. 5c, d), along with the elevated levels of mitotic cells with abnormal spindles (monopolar and multipolar mitotic spindles; Fig. 5c and e). These findings independently show that both these patient-derived CENP-E variants can adversely impact on normal mitotic spindle formation and chromosome segregation, phenotypes consistent with our observations derived from patient LCLs.

LR05-054a2 LCLs exhibit impaired mitotic progression

To examine the proficiency of mitotic progression in patient LCLs in more detail, we performed a *block-release-block* experiment using colcemid coupled with cytochalasin B treatments. Specifically, LCLs were briefly treated with colcemid (4 h) to synchronize a sub-population of cells in mitosis. The colcemid was removed by washing and cells

then released into cytochalasin B, a cytokinesis inhibitor, for 24 h. The frequency of cytochalasin B-induced binucleate cells was compared to untreated cells for each of the WT, PCNT-MOPDii and LR05-054a2 LCLs. The ratio of binucleate cells, that is the number of binucleates following colcemid block and release into cytochalasin B, over the level of binucleates in untreated (Unt) LCLs, reflects the progression through mitosis following release from the colcemid block. We observed a decreased ratio of binucleate cells following this *block-release-block* procedure in PCNT and CENPE (LR05-054a2)-mutated LCLs compared to WT consistent with a delay in progression through mitosis upon release from colcemid (Fig. 6a).

Expression of the D933N and K1355E variants of CENP-E are associated with delayed mitotic progression

Using our Flp-In (Invitrogen) CENPE system in DLD-1 cells engineered to stably co-express RFP-histone H2B to enable chromosome visualization, we employed live cell imaging to determine the kinetics of mitotic progression (Fig. 6b, c). Whilst we found that expression of p.D933N alone or expression of p.K1355E alone, on average, was associated with a slower progression through mitosis compared to WT CENP-E, these delays were not statistically significant (Fig. 6c). The mean mitotic transit time for WT CENP-E was 117 min compared to 175 min for D933N and 147 min for K1355E. But when both variants were engineered *in-cis* into CENP-E, this resulted in a statistically significant delay in mitosis (205 min) compared to WT CENP-E (117 min) (Fig. 6b, c). These findings show that expression of a CENPE cDNA engineered to carry both patient-derived variants (p.D933N and p.K1355E) *in-cis* resulted in a significantly delayed mitosis. These findings are consistent with the *block-release-block* findings using LR05-054a2 LCLs (Fig. 6a).

LR05-054a1 LCLs exhibit a functional ATR-dependent DNA damage response

Seckel syndrome and MOPDii patient-derived LCLs exhibit defects in the ATR-dependent DDR, including impaired UV-induced G2-M cell cycle checkpoint activation and compromised ATR-DDR dependent signaling (Alderton et al. 2004; Griffith et al. 2008; O'Driscoll et al. 2003; Ogi et al. 2012; Qvist et al. 2011). To determine whether LCLs from patient LR05-054a1 exhibited an impaired ATR-dependent DDR, we treated the cells with different forms of radiation: UV to activate the ATR-DDR and IR to activate the ATM-DDR, and then assessed mitotic index (MI). G2-M cell cycle checkpoint activation is indicated by a decrease in MI due to the cell cycle arrest in G2 phase following radiation. Similar to WT

LCLs, LR05-054a1 exhibited a decrease in MI following UV indicative of a functional ATR-dependent checkpoint (Fig. 7a). This is in contrast to LCLs derived from a patient mutated in ATR (ATR-S; ATR-Seckel syndrome) or in PCNT-mutated MOPDii LCLs (PCNT). All patient-derived LCLs demonstrated a drop in MI following IR indicative of a functional ATM-dependent G2-M checkpoint (Fig. 7a).

Following stalling of DNA replication forks with hydroxyurea (HU), 53BP1 forms microscopically detectable foci in an ATR- and CHK1-dependent manner (Sengupta et al. 2004). Seckel syndrome and PCNT-mutated MOPDii patient-derived LCLs exhibit impaired HU-induced 53BP1 foci formation indicative of an impaired ATR-DDR (Griffith et al. 2008; Ogi et al. 2012). Consistent with the normal UV-induced G2-M checkpoint activation in LR05-054a1, these cells were proficient in HU-induced 53BP1 foci formation, in contrast to ATR-S and PCNT-defective MOPDii LCLs (Fig. 7b). Collectively, these results suggest that LR05-054a1 does not exhibit an impaired ATR-DDR.

Discussion

The developing embryonic neuroepithelium is exquisitely sensitive to perturbations in cell cycle kinetics as this system is characterized by a temporally restricted rapid expansion of stem cells (Rakic 1995). It is also sensitive to perturbations in spindle orientation, as alterations in the cleavage plane can disrupt symmetric-asymmetric division, migration and differentiation (Fish et al. 2006). Furthermore, defects in microtubule dynamics and microtubule interacting proteins are associated with a range of neuronal migration disorders (e.g., lissencephaly, pachygyria, polymicrogyria) coupled with microcephaly (Jaglin and Chelly 2009; Poirier et al. 2013).

Collectively, disruption of centrosome-spindle processes can result in an overlapping set of human pathologies ranging from PM, PM with short stature to overt MPD (Al-Dosari et al. 2010; Bond et al. 2002, 2005; Guernsey et al. 2010; Kalay et al. 2011; Sir et al. 2011). Whilst pathogenic defects have been identified in centrosome, microtubule spindle and spindle-associated components, defects in core kinetochore components underlying MPD with cortical malformation have not yet been reported. Recently, a rare homozygous mutation in CASC5 was reported in three consanguineous families from the same isolated geographical region; one of whom was the original Moroccan family used to define the MCPH4 locus (Genin et al. 2012; Jamieson et al. 1999). CASC5 is a component of the KMN complex (KNL1-Mis12 complex-Ndc80 complex), a kinetochore localized network implicated in several functions

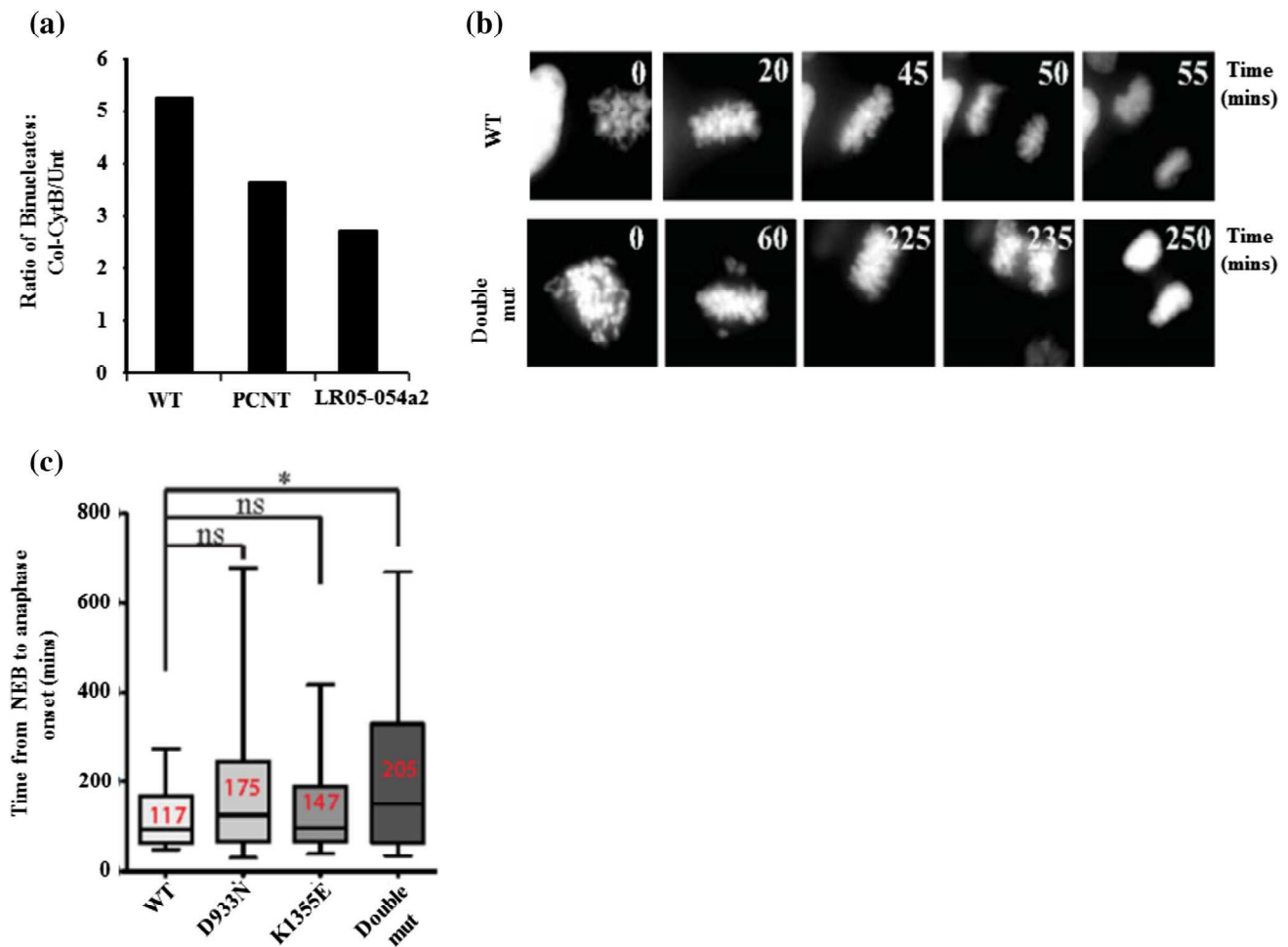


Fig. 6 *CENPE*-mutated patient LCLs exhibit delayed mitotic progression which is also observed upon conditional expression of *CENP-E* carrying D933N and K1355E *in-cis*. **a** The ratio of binucleates following colcemid block and release into cytochalasin B compared to untreated (Unt) cells is lower in *PCNT*-mutated MOPDii (*PCNT*) or *CENPE*-mutated patient derived LCLs (LR05-054a2), compared to WT, consistent with a slower progression through mitosis under these conditions in these patient cells. **b** DLD-1 cells engineered by Flp-In recombination for the conditional (doxycycline-induced) expression of siRNA-resistant *CENPE* and *CENPE* patient variants were plated and 24 h later subjected to *CENPE* siRNA targeting endogenous *CENP-E* only as described in Fig. 5a. These cells also contain a RFP-tagged histone H2B to enable real-time analysis of chromosome dynamics during mitosis by live cell imaging. Approximately 40 h post-siRNA, the cells were treated with doxycycline (Dox; 1 mg/ml) to induce the expression of the siRNA-resistant *CENPE* variants and cells analyzed by live cell imaging. Still images

from the live cell movies following mitosis when wild-type (WT) *CENP-E* was expressed compared to the double mutant (Double mut) expressing D933N and K1355E *in-cis*. The time (in minutes) is indicated in the various panels showing onset of the transition from prophase–metaphase–anaphase–telophase under each conditional expression condition. **c** The time from the onset of nuclear envelope breakdown (NEB) to anaphase was calculated for each *CENP-E* variant when expressed individually or *in-cis* (Double mut; double mutant) and represented in *box-whisker plots*. The average time of the onset of NEB to anaphase is shown in red for each variant. Expression of D933N alone, or K1355E alone, did not result in a statistically significant delay compared to expression of wild-type (WT) *CENPE* (*ns* not significant by Student *t* test), although the average transition for each variant was slightly longer compared to WT. *Expression of both variants *in-cis* did result in a statistically significant delay from NEB to anaphase onset compared to WT ($p < 0.05$ Student *t* test) (color figure online)

including microtubule stabilization and SAC silencing (Bolanos-Garcia et al. 2009; Kiyomitsu et al. 2007, 2011). In contrast to the *CENP-E* patients described here, the *CASC5*-mutated individuals were characterized by PM (age-related OFC ranging from -4 to -7 SD), normal stature, normal motor milestones without evidence of epilepsy or neurological deficits and an otherwise unremarkable

medical history with the oldest patients in their thirties (Genin et al. 2012; Jamieson et al. 1999). Interestingly, detailed analysis *did not* yield any spontaneous mitotic and/or cell cycle defect or evidence of genomic instability in cells from these patients, again in contrast to the *CENP-E* defect described here (Genin et al. 2012). The authors postulated that this may perhaps be due to the restriction of

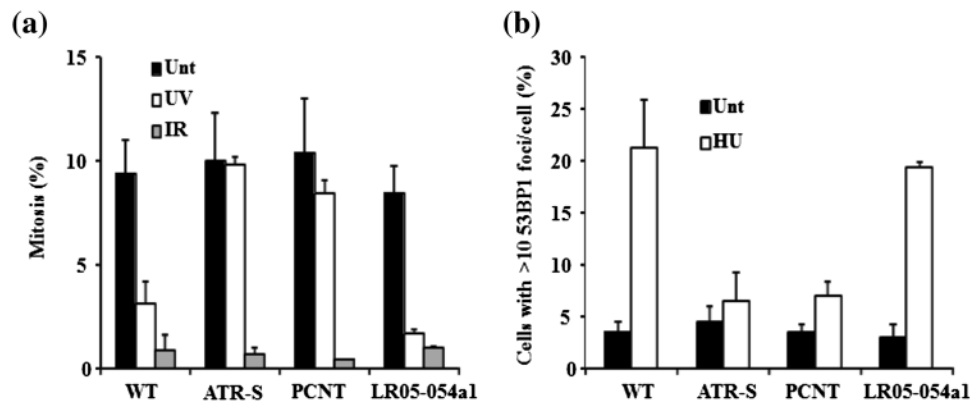


Fig. 7 *CENPE*-mutated patient LCLs exhibit a functional ATR-dependent DNA damage response in contrast to *PCNT*-MOPDii and ATR-Seckel syndrome patient cells. **a** LCLs from wild-type (WT), ATR-mutated Seckel syndrome (ATR-S), *PCNT*-mutated MOPDii (PCNT) and the *CENPE*-mutated male sib (LR05-054a1) were either un-irradiated (Unt), irradiated with ultraviolet light (UV) or ionizing radiation (IR) and their mitotic index (MI % mitosis) enumerated at 4 h. UV-irradiation specifically activates the ATR-dependent G2-M cell cycle checkpoint whilst IR activates the ATM-dependent checkpoint. Both WT and LR05-054a1 LCLs demonstrate a robust

decrease in MI under both conditions. This is in contrast to LCLs from ATR or *PCNT*-mutated patients who exhibit a specific defect in the ATR-mediated G2-M cell cycle checkpoint arrest, as demonstrated previously. **b** Hydroxyurea (HU)-induced 53BP1 foci formation is an ATR and CHK1-dependent process. Cells were either untreated (Unt) or treated with 5 mM HU and 53BP1 foci positive cells enumerated after 2 h via IF. Both WT and LR05-054a1 LCLs demonstrate a similarly increased frequency of cells with 53BP1 foci under these conditions, in contrast to LCLs from ATR or *PCNT*-mutated patients

such anticipated cellular phenotypes to the neuronal system (Genin et al. 2012).

Different defects even within the same centrosomal proteins have previously been found to exhibit a spectrum of growth and neurodevelopmental phenotypes, the basis of which is unclear (Al-Dosari et al. 2010; Bond et al. 2005; Guernsey et al. 2010; Kalay et al. 2011). An important implication of our findings is that hypomorphic defects in other core kinetochore and kinetochore-associated factors (aside from CENP-E) may also underlie MPD in humans. This should be considered when designing targeted sequencing strategies and/or analyzing WES datasets. Our work also illustrates a useful platform to model the functional impacts of candidate gene variants (Flip-In recombination of siRNA-resistant variant carrying cDNA). This is particularly relevant in our situation as one of the patient variant changes reflected the amino acid at the equivalent position in mouse Cenpe. No doubt this variant would likely have been excluded based on standard filtering approaches if used in isolation. Modeling variant pathogenicity in an independent cell system when possible represents a powerful approach when used in combination with patient cell-based functional analysis.

Another important aspect of our findings is that they provide a wider clinical picture for how a hypomorphic defect in a core kinetochore component presents. Nevertheless, with regard to the congenital heart defect (CHD) observed here specifically in LR05-054a1, we cannot be absolutely sure that impaired CENP-E is a causative and/or a contributing factor. Interestingly, CHDs have been reported in

MVA patients, although specifically in those with *CEP57* mutations (*CEP57* encodes a centrosome-associated protein), rather than *BUB1B*. But, the underlying pathomechanism of CHDs in this context has not been described (Snape et al. 2011). As part of our exome analysis of LR05-054a1, we examined 78 genes with known associations with CHDs for potential pathogenic variants but did not find any strong candidates (Online Resource Tables 4 & 5).

The functional analysis of patient LCLs outlined here also demonstrates that defects in spatially distinct proteins that play different roles in mitosis can manifest with similar cellular phenotypes (Online Resource Tables 3). We find that impaired *PCNT* function mimics impaired CENP-E function with respect to CENP-E focalization, CENP-E-dependent BubR1 phosphorylation, mitotic spindle organization and mitotic progression. The basis of this is unclear but may be related to the abnormal spindles characteristic of *PCNT*-MOPDii cells (Rauch et al. 2008). Nevertheless, we also did observe some divergence in cellular phenotypes between CENP-E and *PCNT* (Fig. 4e). Additionally, we did not detect an impaired ATR-dependent DDR in CENP-E LCLs, in contrast to *PCNT*-MOPDii LCLs (Fig. 7a, b). This is likely due to the fact that the centrosome appears to be an important site of localized modification and/or interaction for many DDR factors, including CHK1, a direct substrate of ATR (Kramer et al. 2004; Tibelius et al. 2009; Zhang et al. 2007).

At 2,701 amino acids in length, human CENP-E is one of the largest fibrous extensions emanating from the kinetochore. Structurally, CENP-E is a dimeric kinesin

with a large 230 nm flexible α -helical coiled-coil region (D336-A2471) separating a pair of the head domains that constitute the kinesin motor, ATP binding and microtubule interacting regions (M1-K327), from the C-terminal kinetochore binding domain (Q2472-Q2663) (Fig. 2c) (Kim et al. 2008). Interestingly, defects in other kinesin family proteins with roles in microtubule dynamics and trafficking have been identified in patients with overt cortical structural abnormalities including MIC (Poirier et al. 2013). Both *CENPE* mutations identified here localize to the α -helical coiled-coil region of CENP-E. A noted feature of the structure of CENP-E is the inherent flexibility of the coiled-coil region which is thought to be essential for searching out microtubules (Fig. 2c) (Kim et al. 2008). It is difficult to accurately postulate how together both of the amino acid changes identified here (D933N and K1355E) could impact on CENP-E function. But, the K1355E does represent a change in both charge and size of the side chain (Online Resource Table 2). Our findings provide the basis for further modeling and analysis of these changes at the structural level, as conditional expression of either variant in a model cell system does appear to adversely impact on aspects of normal mitotic progression (Figs. 5, 6).

In summary, we identify rare variants in *CENPE* in two siblings with a profound MPD, structural brain malformations and isolated abnormalities. We catalog various abnormalities in mitotic spindle organization, mitotic progression and segregation in LCLs from these patients, as well as using a model cell system following conditional expression of each *CENPE* variant, alone and together *in-cis*. In demonstrating significant phenotypic overlap between *CENPE*-mutated LCLs and those from a *PCNT*-mutated MOPDii patient, our findings are consistent with a model wherein aberrant mitotic spindle structure, chromosome mis-segregation and delayed mitotic progression represent a significant underlying pathomechanism for MPD in humans. Our findings demonstrate the first example of a core kinetochore defect in contributing to this pathomechanism. This has implications for our understanding of the molecular basis of normal brain and body development and potential for novel genetic defect identification.

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Conflict of interest The authors declare they have no conflict of interest.

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